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1 **Dynamics of particle size on inhalation of environmental aerosol and impact on deposition**
2 **fraction**

3

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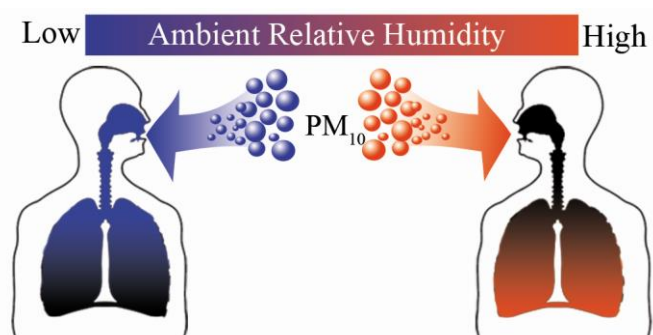
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12

13 Abstract

14 Inhalation of elevated levels of particulate air pollution has been shown to elicit the onset of adverse
15 health effects in humans, where the magnitude of the response is a product of where in the lung the
16 particulate dose is delivered. At any point in time during inhalation the depositional flux of the
17 aerosol is a function of the radius of the droplet, thus a detailed understanding of the rate and
18 magnitude of the mass flux of water to the droplet during inhalation is crucial. In this study, we
19 assess the impact of aerosol hygroscopicity on deposited dose through the inclusion of a detailed
20 treatment of the mass flux of water to account for the dynamics of particle size in a modified version
21 of the standard International Commission on Radiological Protection (ICRP) whole lung deposition
22 model. The ability to account for the role of the relative humidity (RH) of the aerosol prior to, and
23 during, inhalation on the deposition pattern is explored, and found to have a significant effect on the
24 deposition pattern. The model is verified by comparison to previously published measurements, and
25 used to demonstrate that ambient RH affects where in the lung indoor particulate air pollution is
26 delivered.

27 TOC Art



29 1.0 Introduction

30 The association between the inhalation of elevated levels of particulate air pollution and the onset of
31 local and/or systemic adverse health effects in humans has been well established.^{1, 2} The severity of
32 the effect is correlated with the size distribution and chemical composition of the aerosol,³ and overall
33 exposure.⁴ Several inhalation models have been used to predict dose,^{5, 6} typically reporting the
34 deposited fraction of the total aerosol population within a specific region in the lung as a function of
35 initial aerosol diameter. There are numerous challenges in the development of these models that are
36 both biological,⁷ such as accounting for variations in lung morphology and breathing patterns, and
37 physical, such as deposition mechanisms coupled with the fluid dynamics of aerosol transport.⁸ An
38 important factor which remains under-evaluated is the role of the chemical composition of the aerosol

39 particles in influencing the deposition pattern through influencing hygroscopic growth, and the
40 influence of the environmental conditions to which the aerosol is exposed prior to inhalation.^{9,10}

41 The size of aerosol particles in their surrounding environment is dynamic¹¹ and reliant on the relative
42 humidity (RH) in the gas-phase of the aerosol (figure 1A).¹² The absolute magnitude of the size
43 change (the capacity for hygroscopic growth) a single droplet can experience as it drifts from one RH
44 to another is dependent on the chemical composition of the droplet and the two RHs the droplet is
45 transitioning between.¹³ The amount of time taken for the aerosol droplet to reach equilibrium with
46 its surrounding environment, or the rate of change of size, is governed by the kinetics of mass and
47 heat flux to and from the droplet surface (figure 1B).¹⁴ Thus, a detailed understanding of the
48 thermodynamic properties and kinetic processes which control the rate and magnitude of mass
49 transport is required to accurately model aerosol dynamics over the inhalation cycle

50 An aerosol particle will experience an abrupt change in RH during inhalation when the surrounding
51 gas-phase water concentration rapidly increases due to the high humidity in the body. This sudden
52 change in RH will in turn lead to a rapid increase in the diameter of the particle.¹⁵ The RH
53 experienced by the aerosol particle in the environment external to the body, termed the ambient RH,
54 affects the size and composition of the aerosol at the point of inhalation, influencing the capacity of
55 the aerosol to take up further water during inhalation (for example, NaCl in figure 1A). For instance,
56 particles of identical dried solute chemical composition suspended in an atmosphere with an RH of
57 50% (figures 1A and 1C) have greater capacity to absorb water, changing in size by a larger
58 proportion, than those of an identical initial diameter suspended in an atmosphere with a higher RH
59 (figure 1D). Note that the timescale of this size change is dependent on the initial droplet diameter.

60 Lung models fall into one of two broad categories, either whole lung or local scale.⁶ Of these, some
61 are unable to incorporate the hygroscopic condensation that occurs during inhalation¹⁶ while others
62 can,¹⁷ clearly illustrating an absence of consensus in the literature of how the hygroscopic behaviour
63 of inhaled aerosol should be treated. Although local scale models (typically based on simulations
64 using computational fluid dynamics (CFD)) produce more accurate predictions of aerosol deposition
65 within the respiratory tract, validation of these models, *in vivo*, has been impossible until recently.^{18, 19}
66 Instead, local scale deposition has been confirmed using *in vitro* data.²⁰⁻²² In these studies, a portion
67 of the upper respiratory tract is constructed (a plastic nose, mouth and/or throat),²³ aerosol introduced,
68 and then the deposition pattern,²⁴ or total deposition²³ measured. Although highly useful, these *in*
69 *vitro* models are limited to the upper airway and cannot probe in detail the deposition fraction in the
70 deep lung.

71 Though less well defined than CFD models, whole lung models predict regional dose and can be
72 assessed experimentally.^{25, 26} The International Commission on Radiological Protection (ICRP) lung
73 deposition model is considered the standard whole lung model for routine dosimetry assessments.^{27, 28}

74 A model based on the traditional ICRP model is presented here to demonstrate the importance of an
75 accurate parameterization of the hygroscopic properties of the aerosol in predicting the total and
76 regional deposition within the lung and the importance of considering atmospheric conditions when
77 determining aerosol dosage, and to improve the utility of the traditional ICRP model through an
78 accurate parameterization of the hygroscopic growth kinetics during inhalation. Systems in this study
79 range from simple salts to purely organic aerosol. The sensitivity of the regional and total dose to the
80 RH within the lung is also considered. .

81

82 **2.0 Methods**

83 2.1 Model structure

84 The structure of the inhalation model developed here is that of the ICRP model (figure 2A) described
85 in detail previously⁵; sensitivity analysis of the traditional ICRP model has been reported
86 previously.²⁹⁻³¹ Only a brief description of the model will be given, followed by the specific details
87 of the modifications developed here. The respiratory tract is divided into five separate regions: nose
88 (extrathoracic, ET-1), mouth and throat (ET-2), bronchial (BB), bronchiolar (bb) and the alveolar-
89 interstitial (AI) (figure 2A). During a single breath through the nose, the inhalable aerosol passes
90 through ET-1, ET-2, BB and bb twice, once during inhalation and once during exhalation, and the AI
91 only once. The time taken for each breath and the subsequent time taken for the aerosol to reach each
92 region during both inhalation and exhalation are calculated based on many factors, including sex,
93 relative body size and level of activity. The physical mechanism of particle deposition onto the lung
94 surface occurs via diffusion and sedimentation (collectively termed thermodynamic deposition), or
95 impaction (termed aerodynamic deposition). In each region of the lung, the fraction of all aerosol
96 removed from the air, η , is calculated from:

97 Eq. 1 $\eta = 1 - e^{-ar^p}$

98 a and p are constants, and r is a function of the aerodynamic diameter of the aerosol and the air flow
99 rate. The values of a , p and r are unique to each anatomical region and for the mechanism of
100 deposition.³² A table summarizing the previously reported values of a , p and r is given in Supporting
101 Information (Table S1). Both thermodynamic and aerodynamic depositions are highly dependent on

102 the size of the particle. For example, in the AI region of a resting Caucasian male, the aerodynamic
 103 deposition (η_{ae}) is:

104 Eq. 2
$$\eta_{ae} = 1 - e^{(-0.146(2.3d_{ae}^2)^{0.6495})}$$

105 The thermodynamic deposition (η_{th}) is:

106 Eq. 3
$$\eta_{th} = 1 - e^{(-67(2.3D)^{0.6101})}$$

107 d_{ae} is the aerodynamic diameter of the particles and D is the diffusion coefficient of the aerosol (which
 108 itself is a function of the particle diameter). Thus it is critical to have a detailed understanding of the
 109 aerosol dynamics during inhalation to accurately predict deposition frequency. In the standard ICRP
 110 model, the parameterization of hygroscopic growth of the aerosol during inhalation is based on fits to
 111 experimental data³³ and is represented by equation 4:

112 Eq. 4:
$$d_{ae}(t) = (d_{ae}(\infty) \times d_{ae}(0)) - (d_{ae}(\infty) \times d_{ae}(0) - d_{ae}(0)) \times \left(\exp\left(\frac{-(10 \times t)^{0.55}}{d_{ae}(0)}\right) \right)^{0.6}$$

113 The meaning of these variables is given in Table 1. A value is selected for $d_{ae}(\infty)$, typically assumed
 114 to be 3.0 for hygroscopic NaCl droplets.³⁴ Predictions of the time dependent growth of aerosol during
 115 inhalation, using this parameterization for varying values of $d_{ae}(\infty)$ at the specific time points (where
 116 each time point is when the aerosol is in each region in the lung identified in figure 2A), are shown in
 117 figure 2B. Parameters such as the ambient RH prior to inhalation were either not considered or
 118 assumed to be inconsequential (Eq. 4).³³

119 2.2 Semi-analytical treatment for hygroscopic growth kinetics of a single particle

120 In recent work³⁵⁻³⁸ we have performed detailed measurements of the evaporation of water from
 121 numerous aerosol types to verify the semi-analytical solution to the mass and heat flux equations of
 122 Kulmala *et al.*³⁹ These continuum regime equations account for the Kelvin effect and incorporate the
 123 transition corrections factors of Fuchs-Sutugin to broaden their predictive potential into the submicron
 124 regime. The instantaneous mass flux to or from a droplet, I (units g/s), can be calculated from
 125 equation 5:

126 Eq. 5
$$I = -4\pi a \frac{S_\infty - a_w}{\frac{RT_\infty}{M\beta_m AD_w p_{v,\infty}} + \frac{S_a L^2 M}{R\beta_T KT_\infty^2}} \left(\frac{Sh}{2} \right)$$

127 The definition of these variables is given in Table 1. The method by which this equation is used to
 128 simulate the time-dependence in droplet size has been described in detail in previous publications.³⁵
 129 Physically, Eq. 5 describes the continuum diffusional transport of water between the gas phase at
 130 infinite distance and the droplet surface, and the ensuing condensational (or evaporative) kinetics.
 131 Primarily this mass flux is governed by difference between the water activity within the droplet (and,
 132 thus, the droplet vapour pressure and partial pressure of water at the droplet surface) and the degree of
 133 saturation (the partial pressure of water in the gas phase) at infinite distance. In extreme cases,
 134 transport across the droplet surface may be slowed by other kinetic factors such as the presence of a
 135 surfactant coating (factored in through the transitional correction factor β_M) or through the slow
 136 diffusion of water within the bulk of the particle (not considered by equation 5).⁴⁰ Additionally, Eq. 5
 137 accounts for the correction to the mass flux required to correct for the slow heat flux to/from the
 138 particle and the accompanying displacement in droplet temperature away from that of the gas phase.

139 We use equation 5 to predict the time-dependent growth kinetics of aerosol of varying
 140 composition/hygroscopic response during inhalation. The diameter of the aerosol droplets at
 141 appropriate points in time (indicative of the time reached in each region in the lung, figure 2A) were
 142 extracted and used to refine the predictions from the conventional ICRP framework. Then, the
 143 subsequent deposition fraction was estimated. Unless otherwise stated, the assumption is made that
 144 the thermal accommodation coefficient is unity; a range of values of the mass accommodation
 145 coefficient are considered (1.0 to 5.0×10^{-5}).⁴¹ Further, it is assumed that there is no bulk limitation
 146 to mass transfer, i.e. diffusional mixing within the droplet is much more rapid than the timescale of
 147 water transport such that the particles remained homogeneous.¹⁴ The water activity in the droplet at a
 148 particular radial growth factor is critical in determining the flux; the association between water
 149 activity and radial growth factor determines the extent (figure 1A) and rate (figure 1B) at which a
 150 given aerosol will grow following inhalation. For the inorganic aerosols studied here, the radial
 151 growth factor at a corresponding water activity is determined using the Aerosol Diameter Dependent
 152 Model (ADDEM) (Supplemental Figure 1A).^{42, 43} For the organic aerosol, GF is derived using κ -
 153 Kohler theory (Supplemental Figure 1B), according to:

154 Eq. 6
$$GF = \sqrt[3]{1 + \frac{\kappa a_w}{1 - a_w}}$$

155 where κ is the dimensionless hygroscopicity parameter.⁴⁴ For multiple component aerosols, whether
 156 multiple inorganic or organic species (or a mixture of the two), the hygroscopic behaviour of the
 157 aerosol can be readily estimated via the Zdanovskii-Stokes-Robinson (ZSR) approach.⁴⁵⁻⁴⁷

158 A comparison of the predicted growth kinetics of pure NaCl droplets using both the standard ICRP
 159 hygroscopic response and the semi-analytic kinetic model is shown in figure 2B. Differences between

160 the ICRP treatment using the conventional hygroscopic response and the refined treatment are
161 apparent, regardless of the RH in which the aerosol is equilibrated or the initial size. The difference
162 in this size region greater than 50 μm is largely unimportant as the deposition of aerosols within this
163 size regime tends to be 100% of the inhalable fraction.

164 2.3 Conditions of the Model

165 All parameters used for the model, aside from the specific diameter growth factor/water activity
166 relationships for the aerosol considered here, have been described in detail previously.³² Unless
167 otherwise stated, the conditions were those for a healthy Caucasian male breathing through his nose
168 while doing light exercise with an RH within the lung of 99%. The RH in the lung of a healthy
169 individual has been estimated to reach near saturation with an RH between 99% and 99.5%.^{48, 49} The
170 simulations from the model developed here are identified in the figures by the ambient RH
171 immediately prior to inhalation. The simulations produced using the traditional ICRP model are
172 identified by the $d_{ae}(\infty)$ value (Eq. 4).

173 Note that not all particle growth occurring during inhalation may be steady; rather rapid non-linear
174 size changes of crystalline aerosol particles are known to occur through deliquescence (e.g. RH of
175 75% for NaCl) during inhalation. This would result in inhaled dry NaCl particles undergoing no
176 growth in the upper airways followed by rapid growth in the deep lung. To simplify the model, all
177 aerosol is maintained as a liquid, where the solute concentration ranged from dilute, to saturated,
178 through to supersaturated (between 50% RH and 75% for NaCl). Salt particles are only modelled
179 down to an RH of 50% while organic aerosol are modelled down to RH $<20\%$.

180

181 **3.0 Results**

182 3.1 Validation of model through comparison with experimental data

183 Two commonly referenced studies that examined the deposition of mono-disperse aerosol in the
184 respiratory tract were reported by Tu and Knutson,⁵⁰ and by Blanchard and Willeke.⁵¹ The structure
185 of these studies was as follows: NaCl^{50, 51} (or kerosene⁵⁰) was nebulized, dried, size selected and held
186 in a Teflon bag. Immediately prior to inhalation, Tu and Knutson raised the RH of the droplets to
187 95%, while Blanchard and Willeke left the aerosol dry. The difference in aerosol counts between the
188 inhaled and exhaled air was then measured. The results of their studies are shown in figure 3.

189 All of the parameters and experimental conditions used by Tu and Knutson, from the generation of the
190 aerosol, to the pre-treatment and biological factors such as lung volume and sex, are able to be
191 accounted for in the model developed here and a comparison is shown in figure 3. Excellent

192 agreement was found for both the hygroscopic (NaCl) and hydrophobic (kerosene) aerosols. Proper
193 consideration of the pre-treatment of the NaCl aerosol prior to inhalation was essential to accurately
194 predict the deposition pattern (figure 3A).

195 The importance of quantifying the hygroscopicity of the aerosol is shown in figure 3B. Kerosene is a
196 hydrocarbon liquid consisting of a mixture of carbon chains between 6 and 16 carbon per molecule.
197 The κ value of a low solubility organic aerosol is $\sim 0.03^{52}$. Excellent agreement in total dose between
198 the model and experimental results is observed only if the aerosol is assumed to have low
199 hygroscopicity with values between 0.05 and 0.01. This demonstrates that through accurately
200 simulating the dynamics of organic aerosol growth, accurate predictions of aerosol deposition within
201 the lung are possible in the ICRP framework.

202

203 3.2 The ambient RH affects the deposition pattern of pure inorganic aerosol

204 The influence of the RH prior to inhalation on the deposition pattern of NaCl within the lung is shown
205 in figure 4A. Of particular interest in this study is how specific changes in the model conditions
206 affected the deposition pattern. For ease of comparison between multiple deposition patterns, rather
207 than reporting multiple full deposition patterns, we report the differences in the deposition fractions
208 for two different environments. In doing this, the subtle changes in the deposition pattern become
209 immediately apparent (figure 4B). For particles larger than $0.1 \mu\text{m}$ in diameter, lowering the RH prior
210 to inhalation was found to lead to a significant increase in lung deposition in every region, with the
211 largest increases observed in the alveolar-interstitial region and, to a slightly lesser extent, in the
212 bronchial region. Deposition in this size regime is governed by aerosol impaction and sedimentation,
213 suggesting that increasing the magnitude of the size change of particles during inhalation increases the
214 influence of these deposition mechanisms. Below $0.1 \mu\text{m}$, lowering the RH prior to inhalation has the
215 opposite effect, slightly lowering the aerosol deposition in the bronchi and alveolar regions. The
216 reason for this reduction in deposition fraction is attributed to the increase in aerodynamic diameter
217 for the aerosol particles following inhalation: the particles become too large to be removed via
218 diffusion while remaining too small to be removed via sedimentation.

219 Similar comparisons of the deposition pattern of NaCl aerosols, as predicted by the standard ICRP
220 model (with $d_{ae}(\infty) = 3$) and for NaCl aerosol at an ambient RH of 50% (as predicted by our model)
221 are shown in figure 4C. For aerosol larger than 100 nm , a similar deposition pattern was observed for
222 these two simulations; this was not unexpected given that the ICRP model used with a $d_{ae}(\infty) = 3$ is
223 commonly used to predict total and regional dose of NaCl.

224 When the ambient RH is increased to 90% (figure 4D), a more pronounced difference between the
225 traditional ICRP estimation and the refined model is observed in the deposition pattern. Below 100
226 nm, the ICRP underestimates the overall deposition by ~20% due to an overestimation in the
227 magnitude of water uptake when the surface curvature (or Kelvin effect) is not considered (figure 2B).
228 The importance of the Kelvin effect in aerosol dynamics is well understood, hence its integration in
229 numerous lung deposition models.^{53, 54} The absence of an appreciation of the Kelvin effect in the
230 standard ICRP model limits its accuracy for predicting aerosol deposition for this size region. Figure
231 4D clearly demonstrates the limitations of the standard ICRP model: ambient conditions are not
232 considered which leads to inaccurate estimations of dose.

233 Due to the insensitivity of the radial growth dependence of aerosol on RH over the ambient
234 temperature range 0 – 40 °C, changing the ambient temperature was found to have no effect on the
235 deposition pattern (data not shown).

236 The effect of aerosol hygroscopicity on the deposition pattern for aerosol containing similar species
237 was examined. NaCl and ammonium sulphate ((NH₄)₂SO₄) both have well-defined deliquescence
238 (75% for NaCl and 79% for (NH₄)₂SO₄) and efflorescence (45% for NaCl and 37% for (NH₄)₂SO₄)
239 RHs, but show subtle differences in their radial growth factors when the aerosol is liquid
240 (Supplemental Figure 1A). Despite NaCl and (NH₄)₂SO₄ having similar hygroscopic behaviour, a
241 significant difference in the deposition pattern in the lung was observed in their inhalation simulations
242 (Supplemental Figure 2); the largest difference, of 14%, between the two deposition patterns was for
243 aerosol with a diameter around 700 nm.

244

245 3.3 The RH in the lung affects the deposition pattern of pure NaCl aerosol

246 The RH within the alveolar regions of the lung is considered to be near saturation (~99.5%).⁵⁵ In the
247 upper airways, the RH is dependent on many factors including the RH of the inhaled air, breathing
248 rate and the disease state of the individual. RH probes are notoriously inaccurate at RH above 90%
249 with a typical error associated with the probe in excess of ±2%, thus making direct and reliable
250 measurements of the RH in the upper airway impossible. The sensitivity of the deposition pattern of
251 NaCl aerosol to the assumed humidity within the lung is significant (figure 5 and Supporting
252 Information Figure 3), with upwards of 45% more aerosol ranging in size between 800 to 900 nm
253 being deposited in the healthy lung than the diseased lung (figure 5A).

254

255 3.4 The physicochemical properties of organic aerosol affect the lung deposition pattern

256 Organic aerosols are common to urban environments. Of interest to this study are the differences in
257 the deposition pattern in the lung resulting from changes to the physicochemical of the organic aerosol
258 (figure 6).

259 A reduction in the hygroscopicity parameter κ (see Eq. 6) leads to significant changes in both the rate
260 and magnitude of the size change of the aerosol during inhalation across a broad range of initial
261 diameters (figure 6A). This difference in size leads to a significant change in the deposition pattern in
262 the lung (figure 6B and Supporting Figure 4).

263 Common to organic aerosols is the presence of surface active species that can limit the rate of mass
264 flux to and from the droplet. In modelling aerosol mass flux, the likelihood of water uptake is
265 characterised by the mass accommodation coefficient (α), which is equivalent to the fraction of
266 molecular collisions of water with the surface that lead to absorption within the aerosol droplet bulk.
267 For pure water, α is above 0.5⁵⁶ and is reduced as the surface of the droplet becomes covered with
268 surface active species.³⁵ The sensitivity of the deposition pattern to the mass accommodation
269 coefficient was explored (figure 6C). Minimal differences in the deposition pattern were observed
270 until α was reduced to 5×10^{-4} . For reference, this α value is similar to that for the evaporation
271 coefficient for water transport across the interface of a drying aerosol droplet coated in 1-
272 pentadecanol (C₁₅H₃₁OH).

273

274 3.5 Estimating regional and total dose of indoor air pollution as a function of relative humidity

275 Regardless of aerosol type, the majority of fluctuations in regional and total dose as a function of
276 ambient RH were observed for aerosol between 100 nm and 10 μm . The sources of indoor particulate
277 air pollution are limited, resulting in a size regime encapsulated in this region (centred $\sim 1 \mu\text{m}$ in
278 diameter⁵⁷). Based on the size distribution of indoor particulate air pollution measured by Yang *et al.*
279 (Supporting Figure 6),⁵⁷ the changes in regional and total dose were estimated using the modified
280 ICRP model (figure 7).

281 Ambient RH was found to have little effect on the total dose (figure 7D), or the dose delivered to the
282 exothoracic region (figure 7A). However, the hygroscopic response was shown to shift significantly
283 deposition to the alveolar region from the bronchiolar region as the ambient RH increased.

284 Ambient RH may affect the adverse health effects associated with indoor air pollution (figure 7). The
285 location in the lung that the dose is delivered will be affected by ambient RH, which may have a
286 dramatic effect on the downstream biological response. The potential for adverse health effects

287 associated with these shifts in deposition pattern to result from long term exposure will need to be
288 explored further.

289

290 **4.0 Discussion**

291 A novel adaptation to the traditional ICRP model to incorporate a detailed treatment of the kinetic and
292 thermodynamic response of the aerosol during inhalation is presented. The parameters of the model
293 are presented (Tables 1 and SI1). This model can readily predict the total and regional dose for any
294 aerosol where the hygroscopic growth as a function of RH is known; for numerous pure (and
295 multicomponent) aerosols, there are many aerosol thermodynamic models to predict this behaviour
296 available online (e.g. the Extended AIM Aerosol Thermodynamics Model).

297 The condensational growth of a droplet during inhalation is governed both by the hygroscopic
298 capacity of a droplet to grow (a thermodynamic factor) and the kinetics of mass fluxes. Although the
299 largest changes in the deposition pattern results when the capacity for hygroscopic growth is
300 accurately accounted for, kinetic factors do also play an important role with size changes occurring on
301 the same timescale as inhalation/exhalation, particularly if precise estimations of regional and total
302 dose are required (figure 6C).

303 A common observation seen here is that changes to aerosol growth, whether due to the aerosol
304 composition or the RH (either in the lung or ambient), affect the deposition fraction in opposing ways
305 for particles smaller and larger than 100 nm. For example, consider the effect of RH within the lung
306 itself on regional dose. The airway with an RH of 95% (figure 5A) experiences a substantial decrease
307 in the overall aerosol deposition in the size region between 0.1 and 5 μm when compared to the
308 healthy lung while at the same time an increase in total aerosol deposition for aerosol under 100 nm
309 was predicted for the lung with an RH of 95%. Broadly speaking, limitations to aerosol growth lead
310 to an increase in the deposition frequency of aerosol under 100 nm, and a decrease in the deposition
311 frequency of the aerosol between 0.1 and 10 μm . Thus, the consequences of hygroscopic growth
312 during inhalation on the deposition pattern is dependent on aerosol size (whether it is above or below
313 100 nm) and should be treated as such.⁵⁸

314 The importance of the hygroscopic behaviour of inhaled aerosol on the deposition pattern may be
315 amplified for those with diseases of the lung. For example, asthmatics experience airway constriction
316 and difficulty breathing in humid conditions, which is believed to be triggered by airway sensory
317 nerves that are sensitive to temperature changes.⁵⁹ Additionally, the inhalation of submicron
318 particulates also triggers similar asthmatic symptoms.⁶⁰ In section 3.2, aerosol particles under 100 nm
319 in diameter were shown to deposit to a higher degree when the aerosol growth during inhalation was

320 limited at higher ambient RH (figure 4B). Thus, limitations to aerosol growth during inhalation could
321 lead to a larger dose of sub 100 nm in diameter aerosol particles penetrating deep into the lung and
322 may trigger and amplify asthmatic symptoms.

323 The results of figure 5 may be used to gain insight into both the pathogenesis and treatment of lung
324 diseases such as asthma and chronic obstructive pulmonary disease (COPD), where both conditions
325 result from chronic inflammation and are characterized by airway wall thickening.^{61, 62} The
326 thickening of the airway wall leads to an overall reduction of the humidity within the afflicted lungs.⁶³
327 Both the exacerbation and pathogenesis of these conditions have been associated with the inhalation
328 of particulate air pollution with an aerodynamic diameter under 100 nm. As shown in figure 7, a
329 reduction of humidity in the lung leads to an increase in the deposition of aerosols within this size
330 fraction deep in the lung. In the treatment of asthma and COPD, the drug is delivered directly to the
331 lung through the inhalation of a pharmaceutical-containing aerosol. The aerosols generated from
332 pharmaceutical delivery devices (e.g. nebulizers) typically have a log-normal particle size distribution
333 with a mean aerodynamic diameter between 3 and 5 μm . When compared to the humid lung (RH of
334 99.9%), the dry lung (RH of 95%) shows a reduction in pharmaceutical aerosol deposition, meaning
335 the effective dose of these pharmaceuticals may be reduced by up to 50%. There is increasing evidence
336 that drug deposition into the peripheral lung provides an improved clinical treatment compared to
337 large aerosol treatment.⁶⁴ This reduction in dose to the deep lung has the potential to reduce in drug
338 efficacy; further study is required to explore this further. It should be noted that airway remodelling
339 (e.g. gland enlargement, subepithelial fibrosis, epithelial alterations) can be a common feature of some
340 diseases of the lung, such as emphysema and COPD.⁶⁵ These modifications (e.g. breakdown of tissue,
341 mucus-hypersecretion) will clearly have a significant effect on the aerosol deposition rate in the lung
342 as narrower airways will lead to a higher deposition rate via diffusion (remembering that deposition
343 via diffusion is a function of aerosol diameter/hygroscopic growth). An improved understanding of
344 the interplay between aerosol dynamics and the physical structure of the lung and their combined
345 influence on deposition rates is critical for studying both the treatment, and pathogenesis, of lung
346 disease.

347 Although there is an appreciation of the hygroscopic growth of an aerosol during inhalation on total
348 and regional dose, what is continually overlooked are the conditions of the aerosol prior to inhalation,
349 specifically the RH in which it is situated. An example of this is the article by Mitsakou *et al.*⁵⁴ where
350 they developed a whole lung Eulerian model to predict aerosol deposition. In the model, aerosol
351 growth is calculated using the theory described by Mason;⁶⁶ the humidity in which the aerosol
352 originates is not considered and the aerosol growth calculated is that of an originally dry aerosol. This
353 oversight meant that the aerosol growth in their model may be overestimated, which may explain the
354 slight (~5-10%) over estimation in the deposition fraction observed when their model was compared

355 to experimental data; over estimating aerosol growth leads to an increase in the deposition fraction for
356 aerosol 0.01 to 1 μm in diameter (figure 4). The sensitivity to the ambient RH is typically not
357 considered in lung inhalation models; in the literature review made during the preparation of this
358 manuscript, no articles discussing this point were found. We speculate that this oversight is present in
359 numerous models, and consideration of it should serve to improve other current inhalation models.

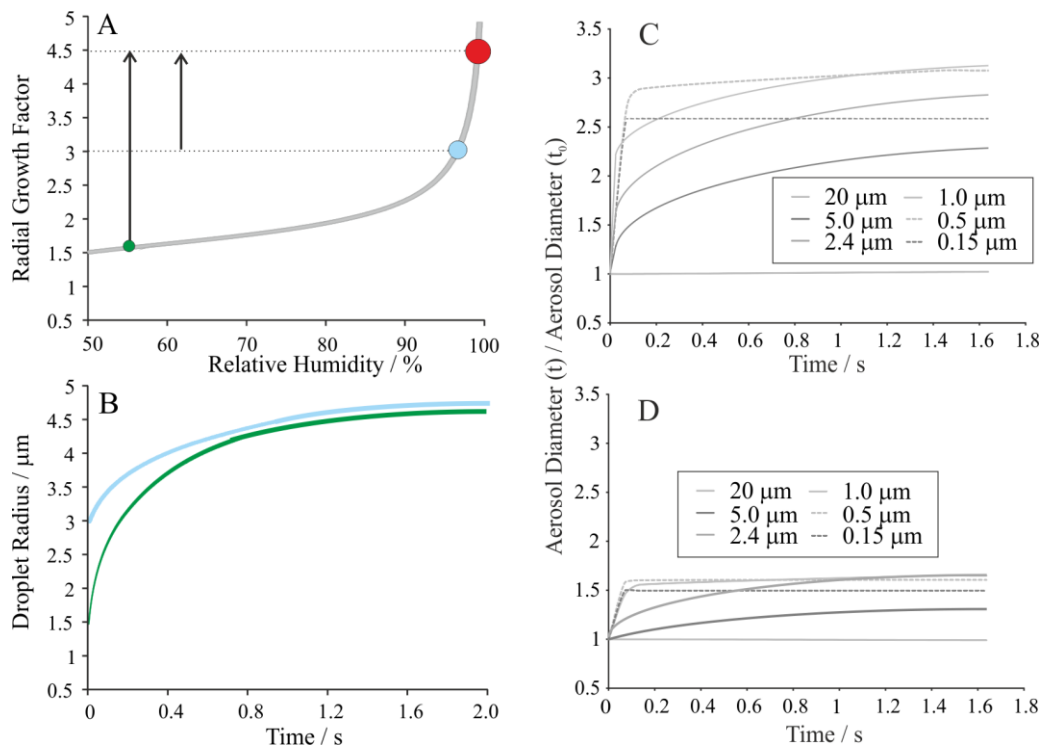
360

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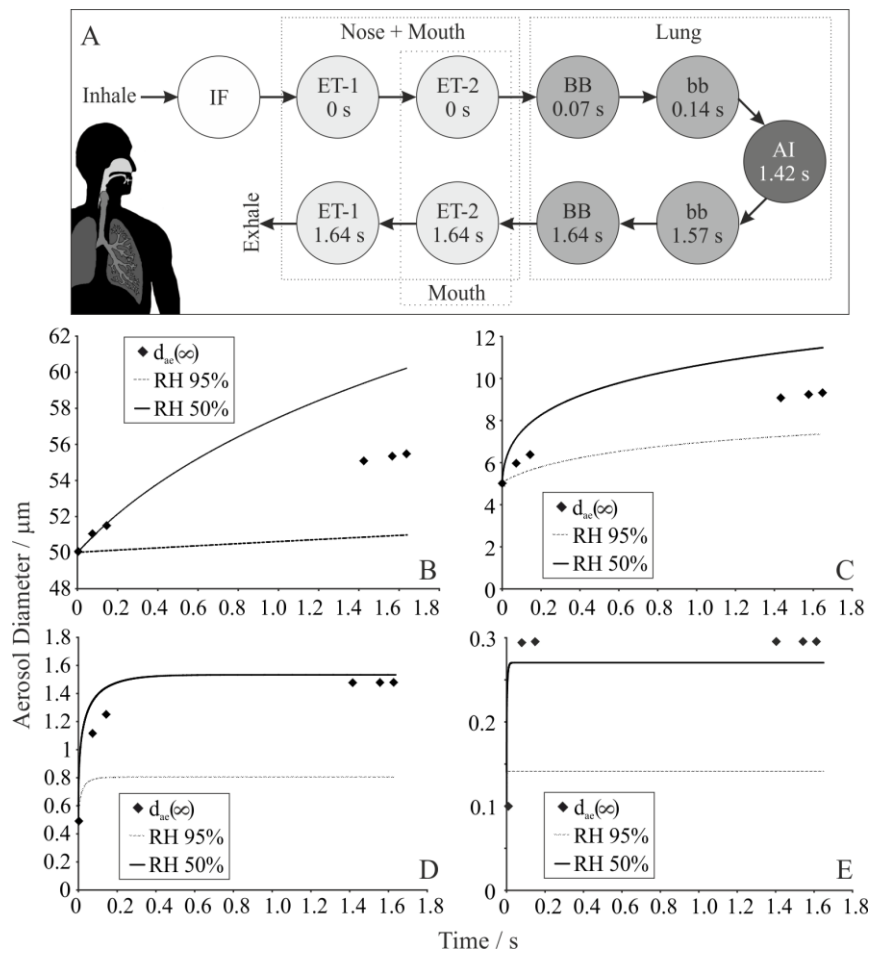
366 Supporting Information is available free of charge via the Internet at <http://pubs.acs.org>. The table
367 and figures include data input in the model (e.g. core equations used to produce the model, radial
368 growth curves, size distributions), and model predictions of regional and total dose as a function of
369 various parameters (e.g. aerosol composition, lung conditions).

370

371 **6.0 Figures**372 **Figure 1**

373

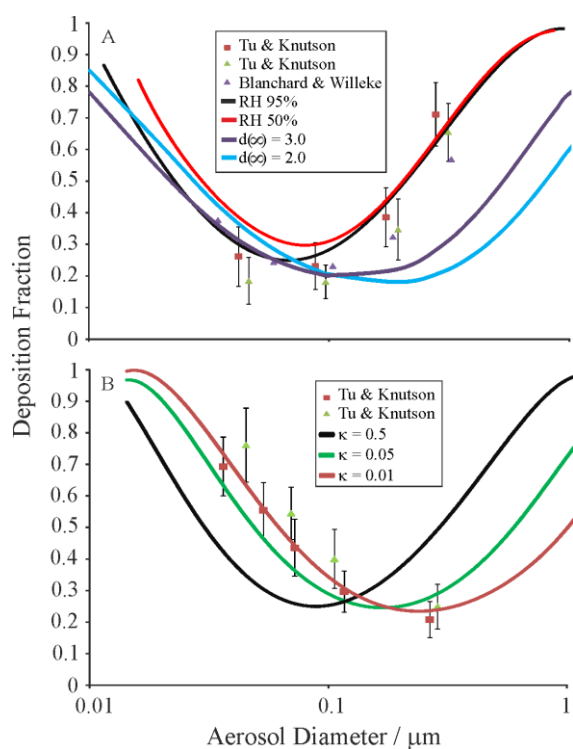
374 The (A) relative magnitude and (B) absolute rate of hygroscopic growth of a single inhaled NaCl
 375 aerosol droplet when the ambient RH prior to inhalation is at 55% (green) and 95% (blue); the dry
 376 radius for both particles is 1 μm . The arrows in (A) show the magnitude of change in radial growth
 377 factor (indicated by the dotted lines) two droplets will experience during inhalation; the droplet
 378 initially suspended in air at RH=55% will grow more than the droplet initially suspended at an
 379 RH=95% during inhalation. The relative growth following inhalation of NaCl aerosols of various
 380 initial diameters that have equilibrated at RHs of (C) 50% and (D) 95%.

381 **Figure 2**

382

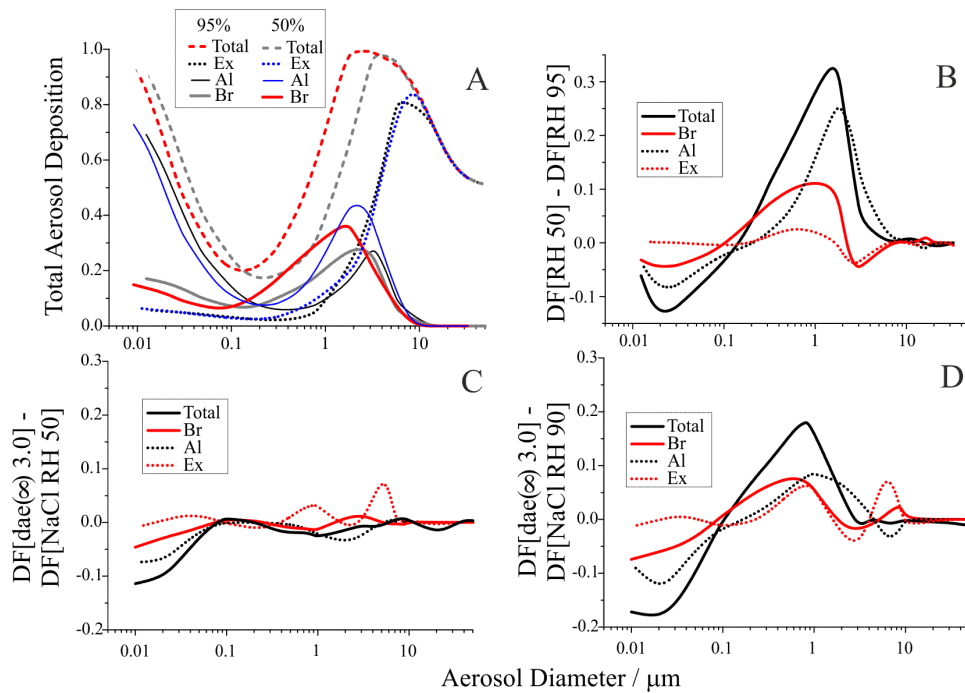
383 (A): The structure of the ICRP model (see text for definitions of regions); each anatomical region of
 384 the lung is indicated by shade for both the inset image and circles. The time point in which the parcel
 385 of air passes through each given region is indicated by number within each circle. The hygroscopic
 386 growth of aerosols of initial diameters of (B) 50 μm , (C) 5 μm , (D) 0.5 μm and (E) 0.05 μm following
 387 inhalation where the RH within the lung is 99% as predicted by the ICRP model ($d_{ae}(\infty) = 3.0$)
 388 (diamonds) and by the semi-analytical solution to the mass and heat flux equations of Kulmala *et al*³⁹
 389 (lines) (where the ambient RH was either 50% or 95%). Note that the time points indicated in the
 390 circles of (A) correspond with those for $d_{ae}(\infty)$ in (B).

391

392 **Figure 3**

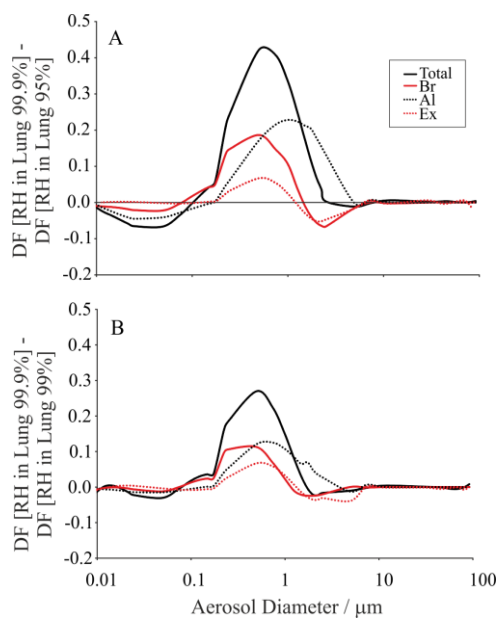
393

394 Comparison of the deposition fraction of (A) NaCl and (B) organic aerosol mimics in the lung
 395 predicted by the model developed here and the traditional ICRP model with the experimental data
 396 collected by Tu & Knutson⁵⁰ and Blanchard & Willeke.⁵¹ For both (A) and (B), the symbols indicate
 397 previously reported experimental data of human exposure. For (A), the lines indicate dose predictions
 398 from the traditional ICRP model ($d(\infty)=3.0$ and $d(\infty)=2.0$) and from the adapted ICRP model reported
 399 here (where the ambient RH was either 50% or 95%). For (B), the lines indicate dose predictions
 400 from only the adapted ICRP model reported here while the hygroscopic behaviour of the aerosol was
 401 estimated by setting the κ -value given into equation 6 (Supplementary Information Figure S-1).

402 **Figure 4**

403

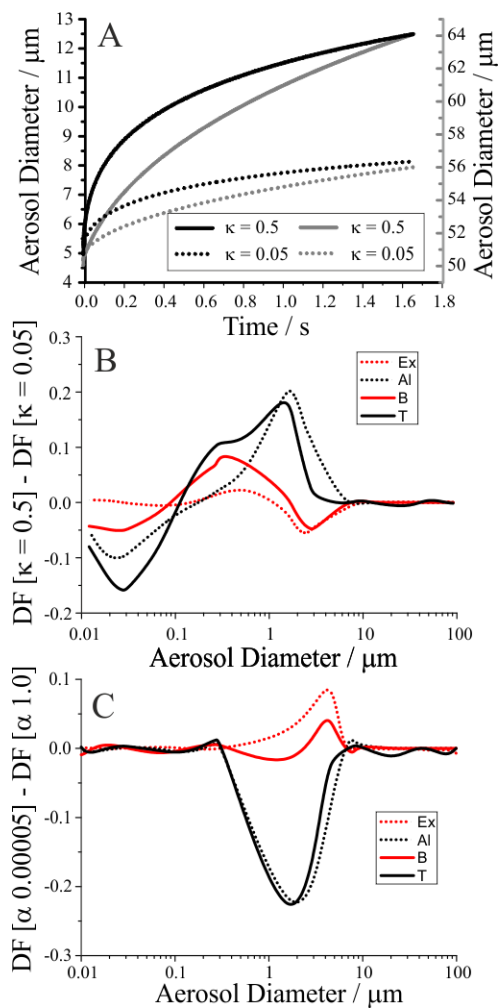
404 (A) Deposition fraction (DF) of NaCl aerosols in the extrathoracic region (Ex), bronchial region (Br),
 405 alveolar-interstitial region (Al) and total lung (Total) as a function of relative humidity (RH) the
 406 aerosol experiences prior to inhalation. Differences in the DF of inhaled aerosol as a function of the
 407 magnitude of aerosol growth during inhalation are also reported. Specifically, changes in the DF of
 408 NaCl aerosols resulting from the conditions of the aerosol immediately prior to inhalation are
 409 explored. (B) is the difference in deposition fraction for NaCl suspended in an ambient air with an
 410 RH of 50% and 95% prior to inhalation. The difference in deposition fraction between the ICRP
 411 model prediction of hydrophilic aerosol and the prediction for NaCl suspended in an ambient air
 412 with an RH of (C) 50% and (D) 95% prior to inhalation.

413 **Figure 5**

414

415

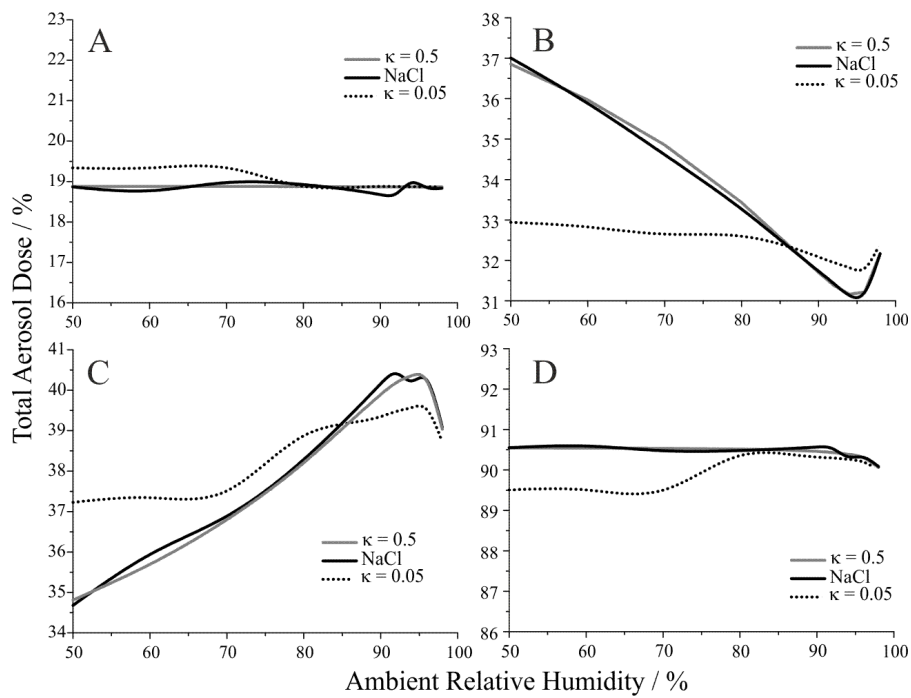
416 Differences in the deposition fraction (DF) of inhaled aerosol (initially at an ambient RH of 50%)
 417 resulting from the changes in RH in the lung as a function of aerosol diameter. Deposition in a lung
 418 with an RH of 99.9% is compared with deposition in a lung with an RH of (A) 95% and (B) 99%.
 419 Total deposition (Total), combined bronchi and bronchiolar (Br), extrathoracic (nose, mouth and
 420 throat) (Ex), and alveolar-interstitial (Al).

421 **Figure 6**

422

423

424 How the growth dynamics of organic aerosol affect total and regional dose. (A) The growth kinetics
 425 of two organic aerosols of different hydrophobicities in the lung where the ambient RH is 20%. The
 426 colour of the line indicates the starting diameter of the droplet (black for 5 micrometers, left y-axis.
 427 And grey for 50 micrometers, right axis) while the line style indicates hygroscopicity of the droplet
 428 (i.e. the value of κ). (B) How this difference in dynamic behaviour of the aerosol described in (A) will
 429 affect the subsequent deposition fraction (DF) of the aerosol in the lung. (C) Surface active species
 430 affect the total and regional doses; the deposition pattern of organic aerosol with a mass
 431 accommodation coefficient (α) of 0.00005 is compared to that of water. Total deposition (T),
 432 combined bronchi and bronchiolar (B), extrathoracic (nose, mouth and throat) (Ex), and alveolar-
 433 interstitial (Al).

434 **Figure 7**

435

436 The percentage of the total dose of indoor particulate air pollution delivered to the exothoracic region
 437 (A), bronchi region (B), alveolar region (C) and the whole (D) lung as a function of ambient RH and
 438 particle type. The specific particle types shown consist of a solute of either pure NaCl or an organic
 439 aerosol mimic for which the hygroscopic behaviour is determined by the value of κ (equation 6), with
 440 a value of 0.5 representing a hygroscopic aerosol and a value of 0.05 an aerosol of low hygroscopic
 441 response. Doses were modelled for a male at rest breathing through his mouth.

442 **7.0 Tables****Table 1: Variables Used in Modified ICRP Model**

Symbol	Unit	Name and/or Meaning
d_{ae}	μm	Aerodynamic diameter
$d_{ae}(0)$	μm	Aerodynamic diameter at the point of inhalation
$d_{ae}(t)$	μm	Aerodynamic diameter at time t
$d_{ae}(\infty)$	Unitless	Equilibrium growth value/Relative growth in the lung the aerosol will reach given an infinite amount of time
t	s	Time from the point aerosol enters respiratory tract
I	g/s	Instantaneous mass flux to or from a droplet
a	μm	Droplet radius
S_{∞}	Unitless	Water saturation ratio in the gas phase of the aerosol
β_m	Unitless	Transitional correction factors for mass
β_T	Unitless	Transitional correction factors for temperature
a_w	Unitless	Activity of water at the surface of the droplet
M	g/mol	Molar mass of water
D_w	cm^2/s	Diffusion coefficient of water in the gas phase
$p_{v,\infty}$	Pa	Saturation vapour pressure of water
A	Unitless	Stefan flow correction
K	W/(mK)	Thermal conductivity of the gas phase
R	$\text{m}^3\text{PaK}^{-1}\text{mol}^{-1}$	Molar gas constant
L	J/kg	Latent heat of vaporization per unit weight of water
Sh	Unitless	Sherwood number/ Accounts for the enhancement of mass flux in an airflow
GF	Unitless	Radial growth factor
κ	Unitless	Dimensionless hygroscopicity parameter from κ -Kohler theory
S_a	Unitless	Activity at the surface of the droplet
T_{∞}	K	Gas temperature
D	cm^2/s	Diffusion coefficient of aerosol

444 **8.0 References**

- 445 1. Pope, C. A.; Thun, M. J.; Namboodiri, M. M.; Dockery, D. W.; Evans, J. S.; Speizer, F. E.;
446 Heath, C. W., Particulate Air-Pollution as a Predictor of Mortality in a Prospective-Study of Us
447 Adults. *Am. J. Respir. Crit. Care Med.* **1995**, *151*, (3), 669-674.
- 448 2. Pope, C. A.; Ezzati, M.; Dockery, D. W., Fine-Particulate Air Pollution and Life Expectancy
449 in the United States. *New Engl J Med* **2009**, *360*, (4), 376-386.
- 450 3. Araujo, J. A.; Nel, A. E., Particulate matter and atherosclerosis: role of particle size,
451 composition and oxidative stress. *Part Fibre Toxicol* **2009**, *6*, (24), 1-19.
- 452 4. Anderson, J. O.; Thundiyil, J. G.; Stolbach, A., Clearing the air: a review of the effects of
453 particulate matter air pollution on human health. *J. Med. Toxicol.* **2012**, *8*, (2), 166-75.
- 454 5. Hofmann, W., Modelling inhaled particle deposition in the human lung-A review. *J Aerosol*
455 *Sci* **2011**, *42*, (10), 693-724.
- 456 6. Rostami, A. A., Computational modeling of aerosol deposition in respiratory tract: a review.
457 *Inhal Toxicol* **2009**, *21*, (4), 262-90.
- 458 7. Yeh, H. C.; Schum, G. M., Models of human lung airways and their application to inhaled
459 particle deposition. *Bull Math Biol* **1980**, *42*, (3), 461-80.
- 460 8. Butler, J. P.; Tsuda, A., Effect of convective stretching and folding on aerosol mixing deep in
461 the lung, assessed by approximate entropy. *J Appl Physiol* **1997**, *83*, (3), 800-9.
- 462 9. Finlay, W. H.; Smaldone, G. C., Hygroscopic behavior of nebulized aerosols: Not as
463 important as we thought? *J Aerosol Med* **1998**, *11*, (4), 193-195.
- 464 10. Finlay, W. H., Estimating the type of hygroscopic behavior exhibited by aqueous droplets. *J*
465 *Aerosol Med* **1998**, *11*, (4), 221-9.
- 466 11. Davies, J. F.; Haddrell, A. E.; Reid, J. P., Time-Resolved Measurements of the Evaporation of
467 Volatile Components from Single Aerosol Droplets. *Aerosol Sci Tech* **2012**, *46*, (6), 666-677.
- 468 12. Chan, C. K.; Liang, Z.; Zheng, J. A.; Clegg, S. L.; Brimblecombe, P., Thermodynamic
469 properties of aqueous aerosols to high supersaturation .1. Measurements of water activity of the
470 system Na⁺-Cl⁻-NO₃⁻-SO₄²⁻-H₂O at similar to 298.15 K. *Aerosol Sci Tech* **1997**, *27*, (3), 324-344.
- 471 13. Clegg, S. L.; Seinfeld, J. H.; Brimblecombe, P., Thermodynamic modelling of aqueous
472 aerosols containing electrolytes and dissolved organic compounds. *J Aerosol Sci* **2001**, *32*, (6), 713-
473 738.
- 474 14. Davies, J. F.; Haddrell, A. E.; Miles, R. E.; Bull, C. R.; Reid, J. P., Bulk, surface, and gas-
475 phase limited water transport in aerosol. *J Phys Chem A* **2012**, *116*, (45), 10987-98.
- 476 15. Broday, D. M.; Georgopoulos, P. G., Growth and deposition of hygroscopic particulate matter
477 in the human lungs. *Aerosol Sci Tech* **2001**, *34*, (1), 144-159.
- 478 16. Khajeh-Hosseini-Dalasm, N.; Longest, P. W., Deposition of particles in the alveolar airways:
479 Inhalation and breath-hold with pharmaceutical aerosols. *J Aerosol Sci* **2015**, *79*, 15-30.
- 480 17. Winkler-Heil, R.; Ferron, G.; Hofmann, W., Calculation of hygroscopic particle deposition in
481 the human lung. *Inhal Toxicol* **2014**, *26*, (3), 193-206.
- 482 18. Conway, J., Lung imaging - two dimensional gamma scintigraphy, SPECT, CT and PET. *Adv*
483 *Drug Deliv Rev* **2012**, *64*, (4), 357-68.
- 484 19. Fleming, J.; Conway, J.; Majoral, C.; Bennett, M.; Caillibotte, G.; Montesantos, S.; Katz, I., A
485 Technique for Determination of Lung Outline and Regional Lung Air Volume Distribution from
486 Computed Tomography. *J Aerosol Med* **2013**, *27*(1), 35-42.
- 487 20. Longest, P. W.; Hindle, M., CFD simulations of enhanced condensational growth (ECG)
488 applied to respiratory drug delivery with comparisons to in vitro data. *J Aerosol Sci* **2010**, *41*, (8),
489 805-820.
- 490 21. Hindle, M.; Longest, P. W., Evaluation of Enhanced Condensational Growth (ECG) for
491 Controlled Respiratory Drug Delivery in a Mouth-Throat and Upper Tracheobronchial Model. *Pharm.*
492 *Res.* **2010**, *27*, (9), 1800-1811.
- 493 22. Byron, P. R.; Hindle, M.; Lange, C. F.; Longest, P. W.; McRobbie, D.; Oldham, M. J.;
494 Olsson, B.; Thiel, C. G.; Wachtel, H.; Finlay, W. H., In Vivo-In Vitro Correlations: Predicting
495 Pulmonary Drug Deposition from Pharmaceutical Aerosols. *J. Aerosol Med. Pulm. Drug Deliv.* **2010**,
496 *23*, S59-S69.

- 497 23. Grgic, B.; Martin, A. R.; Finlay, W. H., The effect of unsteady flow rate increase on in vitro
498 mouth-throat deposition of inhaled boluses. *J Aerosol Sci* **2006**, *37*, (10), 1222-1233.
- 499 24. Holbrook, L. T.; Longest, P. W., Validating CFD predictions of highly localized aerosol
500 deposition in airway models: In vitro data and effects of surface properties. *J Aerosol Sci* **2013**, *59*, 6-
501 21.
- 502 25. Camner, P.; Anderson, M.; Philipson, K.; Bailey, A.; Hashish, A.; Jarvis, N.; Bailey, M.;
503 Svartengren, M., Human bronchiolar deposition and retention of 6-, 8- and 10-micrograms particles.
504 *Exp Lung Res* **1997**, *23*, (6), 517-35.
- 505 26. Davesne, E.; Paquet, F.; Ansoborlo, E.; Blanchardon, E., Absorption of plutonium
506 compounds in the respiratory tract. *J Radiol Prot* **2010**, *30*, (1), 5-21.
- 507 27. Bair, W. J., The Icrp Human Respiratory-Tract Model for Radiological Protection. *Radiat*
508 *Prot Dosim* **1995**, *60*, (4), 307-310.
- 509 28. Bair, W. J., The Revised International Commission on Radiological Protection (Icrp)
510 Dosimetric Model for the Human Respiratory Tract - an Overview. *Inhaled Particles Vii* **1994**, 251-
511 256.
- 512 29. Huston, T. E.; Farfan, E. B.; Bolch, W. E.; Bolch, W. E., Influences of parameter
513 uncertainties within the ICRP-66 respiratory tract model: A parameter sensitivity analysis. *Health*
514 *Phys* **2003**, *85*, (5), 553-566.
- 515 30. Farfan, E. B.; Huston, T. E.; Bolch, W. E.; Han, E.; Beharry, K.; Jupiter, K., Influence of age,
516 gender, and exertion level on dose uncertainties associated with inhalation of weapons-grade
517 plutonium oxide. *Health Phys* **2003**, *84*, (6), S171-S171.
- 518 31. Bolch, W. E.; Huston, T. E.; Farfan, E. B.; Vernetson, W. G.; Bolch, W. E., Influences of
519 parameter uncertainties within the ICRP-66 respiratory tract model: Particle clearance. *Health Phys*
520 **2003**, *84*, (4), 421-435.
- 521 32. Yu, K. N.; Lau, B. M. F.; Nikezic, D., Assessment of environmental radon hazard using
522 human respiratory tract models. *J Hazard Mater* **2006**, *132*, (1), 98-110.
- 523 33. Human respiratory tract model for radiological protection. A report of a Task Group of the
524 International Commission on Radiological Protection. *Annals of the ICRP* **1994**, *24*, (1-3), 1-482.
- 525 34. Londahl, J.; Massling, A.; Swietlicki, E.; Brauner, E. V.; Ketzel, M.; Pagels, J.; Loft, S.,
526 Experimentally Determined Human Respiratory Tract Deposition of Airborne Particles at a Busy
527 Street. *Environ. Sci. Technol.* **2009**, *43*, (13), 4659-4664.
- 528 35. Davies, J. F.; Miles, R. E. H.; Haddrell, A. E.; Reid, J. P., Influence of organic films on the
529 evaporation and condensation of water in aerosol. *P Natl Acad Sci USA* **2013**, *110*, (22), 8807-8812.
- 530 36. Davies, J. F.; Haddrell, A. E.; Rickards, A. M. J.; Reid, J. P., Simultaneous Analysis of the
531 Equilibrium Hygroscopicity and Water Transport Kinetics of Liquid Aerosol. *Anal Chem* **2013**, *85*,
532 (12), 5819-5826.
- 533 37. Haddrell, A. E.; Hargreaves, G.; Davies, J. F.; Reid, J. P., Control over hygroscopic growth of
534 saline aqueous aerosol using Pluronic polymer additives. *Int J Pharm* **2013**, *443*, (1-2), 183-192.
- 535 38. Haddrell, A. E.; Davies, J. F.; Miles, R. E. H.; Reid, J. P.; Dailey, L. A.; Murnane, D.,
536 Dynamics of aerosol size during inhalation: Hygroscopic growth of commercial nebulizer
537 formulations. *Int. J. Pharm.* **2014**, *463*, (1), 50-61.
- 538 39. Kulmala, M.; Vesala, T.; Wagner, P. E., An Analytical Expression For the Rate of Binary
539 Condensational Particle Growth. *Proc R Soc A* **1993**, *441*, (1913), 589-605.
- 540 40. Bones, D. L.; Reid, J. P.; Lienhard, D. M.; Krieger, U. K., Comparing the mechanism of
541 water condensation and evaporation in glassy aerosol. *P Natl Acad Sci USA* **2012**, *109*, (29), 11613-
542 11618.
- 543 41. Miles, R. E. H.; Knox, K. J.; Reid, J. P.; Laurain, A. M. C.; Mitchem, L., Measurements of
544 Mass and Heat Transfer at a Liquid Water Surface during Condensation or Evaporation of a
545 Subnanometer Thickness Layer of Water. *Phys Rev Lett* **2010**, *105*, (11), 116101(1)-116101(4).
- 546 42. Topping, D. O.; McFiggans, G. B.; Coe, H., A curved multi-component aerosol
547 hygroscopicity model framework: Part 1 - Inorganic compounds. *Atmos Chem Phys* **2005**, *5*, 1205-
548 1222.
- 549 43. Topping, D. O.; McFiggans, G. B.; Coe, H., A curved multi-component aerosol
550 hygroscopicity model framework: Part 2 - Including organic compounds. *Atmos Chem Phys* **2005**, *5*,
551 1223-1242.

- 552 44. Petters, M. D.; Kreidenweis, S. M., A single parameter representation of hygroscopic growth
553 and cloud condensation nucleus activity. *Atmos Chem Phys* **2007**, *7*, (8), 1961-1971.
- 554 45. Zdanovskii, A. B., Novyi Metod Rascheta Rastvorimostei Elektrolitov V
555 Mnogokomponentnykh Sistemakh .2. *Zh Fiz Khim+* **1948**, *22*, (12), 1486-1495.
- 556 46. Zdanovskii, A. B., Novyi Metod Rascheta Rastvorimostei Elektrolitov V
557 Mnogokomponentnykh Sistemakh .1. *Zh Fiz Khim+* **1948**, *22*, (12), 1478-1485.
- 558 47. Stokes, R. H.; Robinson, R. A., Interactions in Aqueous Nonelectrolyte Solutions .I. Solute-
559 Solvent Equilibria. *J Phys Chem-Us* **1966**, *70*, (7), 2126-&.
- 560 48. Ferron, G. A.; Haider, B.; Kreyling, W. G., A Method for the Approximation of the Relative-
561 Humidity in the Upper Human Airways. *Bull. Math. Biol.* **1985**, *47*, (4), 565-589.
- 562 49. Cothorn, C. R.; Smith, J. E., *Environmental radon*. Plenum Press: New York, 1987; p xiii,
563 363 p.
- 564 50. Tu, K. W.; Knutson, E. O., Total Deposition of Ultrafine Hydrophobic and Hygroscopic
565 Aerosols in the Human Respiratory System. *Aerosol Sci Tech* **1984**, *3*, (4), 453-465.
- 566 51. Blanchard, J. D.; Willeke, K., Total deposition of ultrafine sodium chloride particles in human
567 lungs. *J Appl Physiol* **1984**, *57*, (6), 1850-6.
- 568 52. Carrico, C. M.; Petters, M. D.; Kreidenweis, S. M.; Sullivan, A. P.; McMeeking, G. R.; Levin,
569 E. J. T.; Engling, G.; Malm, W. C.; Collett, J. L., Water uptake and chemical composition of fresh
570 aerosols generated in open burning of biomass. *Atmos Chem Phys* **2010**, *10*, (11), 5165-5178.
- 571 53. Kleinstreuer, C.; Feng, Y., Lung Deposition Analyses of Inhaled Toxic Aerosols in
572 Conventional and Less Harmful Cigarette Smoke: A Review. *Int J Env Res Pub He* **2013**, *10*, (9),
573 4454-4485.
- 574 54. Mitsakou, C.; Helmis, C.; Housiadas, C., Eulerian modelling of lung deposition with sectional
575 representation of aerosol dynamics. *J Aerosol Sci* **2005**, *36*, (1), 75-94.
- 576 55. Labiris, N. R.; Dolovich, M. B., Pulmonary drug delivery. Part I: Physiological factors
577 affecting therapeutic effectiveness of aerosolized medications. *Brit J Clin Pharmacol* **2003**, *56*, (6),
578 588-599.
- 579 56. Miles, R. E. H.; Reid, J. P.; Riipinen, I., Comparison of Approaches for Measuring the Mass
580 Accommodation Coefficient for the Condensation of Water and Sensitivities to Uncertainties in
581 Thermophysical Properties. *J. Phys. Chem. A* **2012**, *116*, (44), 10810-10825.
- 582 57. Yang, F. X.; Ding, J. J.; Huang, W.; Xie, W.; Liu, W. P., Particle Size-Specific Distributions
583 and Preliminary Exposure Assessments of Organophosphate Flame Retardants in Office Air
584 Particulate Matter. *Environ. Sci. Technol.* **2014**, *48*, (1), 63-70.
- 585 58. Martonen, T. B.; Barnett, A. E.; Miller, F. J., Ambient Sulfate Aerosol Deposition in Man -
586 Modeling the Influence of Hygroscopicity. *Environ Health Persp* **1985**, *63*, (Nov), 11-24.
- 587 59. Hayes, D., Jr.; Collins, P. B.; Khosravi, M.; Lin, R. L.; Lee, L. Y., Bronchoconstriction
588 triggered by breathing hot humid air in patients with asthma: role of cholinergic reflex. *Am J Respir*
589 *Crit Care Med* **2012**, *185*, (11), 1190-6.
- 590 60. Donaldson, K.; Gilmour, M. I.; MacNee, W., Asthma and PM10. *Respir Res* **2000**, *1*, (1), 12-
591 5.
- 592 61. Halwani, R.; Al-Muhsen, S.; Hamid, Q., Airway remodeling in asthma. *Curr. Opin.*
593 *Pharmacol.* **2010**, *10*, (3), 236-45.
- 594 62. Hogg, J. C.; Timens, W., The pathology of chronic obstructive pulmonary disease. *Annu Rev*
595 *Pathol* **2009**, *4*, 435-59.
- 596 63. Matsumoto, H.; Niimi, A.; Tabuena, R. P.; Takemura, M.; Ueda, T.; Yamaguchi, M.;
597 Matsuoka, H.; Jinnai, M.; Chin, K. Z.; Mishima, M., Airway wall thickening in patients with cough
598 variant asthma and nonasthmatic chronic cough. *Chest* **2007**, *131*, (4), 1042-1049.
- 599 64. van den Berge, M.; ten Hacken, N. H.; van der Wiel, E.; Postma, D. S., Treatment of the
600 bronchial tree from beginning to end: targeting small airway inflammation in asthma. *Allergy* **2013**,
601 *68*, (1), 16-26.
- 602 65. Hogg, J. C.; Chu, F.; Utokaparch, S.; Woods, R.; Elliott, W. M.; Buzatu, L.; Cherniack, R.
603 M.; Rogers, R. M.; Sciurba, F. C.; Coxson, H. O.; Pare, P. D., The nature of small-airway obstruction
604 in chronic obstructive pulmonary disease. *New Engl J Med* **2004**, *350*, (26), 2645-2653.
- 605 66. Mason, J., Physics of Cloud and Precipitation. *B Am Meteorol Soc* **1971**, *52*, (12), 1209-&.